Amendment to the Claims:

This listing of claims will replace all prior versions, and listings, of claims in the application:

Listing of Claims:

1. (Original) A method for diagnosis of a disorder associated with the development of beta

amyloid deposits or fibrils in a human or animal subject or assessing the efficacy of treatment

rendered to the subject for such disorder, said method comprising the step of:

A) determining the presence of mtDNA CR mutations.

2. (Original) A method according to Claim 1, wherein Step A comprises making a

qualitative determination that mtDNS CR mutation is or is not present.

3. (Original) A method according to Claim 1, wherein Step A comprises making a

quantitative determination of mtDNS CR mutations.

4. (Original) A method according to Claim 3 further comprising the step of:

B) comparing a mtDNS CR value obtained by the quantitative determination made in

Step A with a control mtDNS CR value to determine whether the subject has significantly more

mtDNS CR mutations than control.

5. (Original) A method according to Claim 3 further comprising the step of:

B) comparing a mtDNS CR value obtained by the quantitative determination made in

Step A with a mtDNS CR value representative of subjects who suffer from a disorder associated

with the development of beta amyloid deposits or fibrils.

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- (Original) A method according to any of Claim 1 wherein Step A comprises testing for a T4141G mutation.
- (Original) A method according to any of Claim 1 wherein Step A comprises testing for a T414C mutation.
- 8. (Original) A method according to any of Claim 1 wherein Step A comprises testing for a T477C mutation
- (Original) A method according to any of Claim 1 wherein Step A comprises testing for a T146C mutation.
- (Original) A method according to any of Claim 1 wherein Step A comprises testing for a T152C mutation.
- (Original) A method according to any of Claim 1 wherein Step A comprises testing for a A189G mutation.
- (Original) A method according to any of Claim 1 wherein Step A comprises testing for a T195C mutation.
- (Original) A method according to Claim 1 wherein Step A is carried out at least in part by PNA-clamping PCR.
- 14. (Original) A method according to Claim 1 wherein Step A is carried out at least in part by oligonucleotide hybridization.
- 15. (Original) A method according to Claim 1 wherein Step A is carried out at least in part by primer extension.

16. (Original) A method according to Claim 1 wherein Step A is carried out at least in part by restriction digestion.

17. (Original) A method according to Claim 1 wherein the determination of Step A is made in a specimen of tissue, cells or body fluid selected from the group consisting of:

i. brain tissue;

ii. brain tissue from the frontal cortex:

iii. nervous tissue:

iv. nerve cells

v. blood

vi. blood cells;

vii. urine;

viii. urinary tract cells;

ix. skin:

x. skin cells:

xi. epithelium:

xii. epithelial cells;

xiii. fibroblasts;

xiv. cerebrospinal fluid; and

xv. cells contained in cerebrospinal fluid.

18. (Original) A method according to Claim 1 wherein the method is carried out for post-symptomatic diagnosis of a disorder in a subject who has begun to exhibit symptoms of that disorder.

19. (Original) A method according to Claim 1 wherein the method is carried out for presymptomatic diagnosis of a disorder in a subject who has not begun to exhibit symptoms of that disorder.

- (Original) A method according to Claim 1 wherein the disorder is a neurodegenerative disease.
- 21. (Original) A method according to Claim 1 wherein the disorder is Alzheimer's Disease.
- 22. (Original) A method according to Claim 1 wherein the disorder is Parkinson's Disease.
- (Original) A method according to Claim 1 wherein the disorder is Down's Syndromeassociated dementia.
- (Original) A method according to Claim 1 wherein the disorder is a spongiform encephalopathy.
- (Original) A method according to Claim 1 wherein the disorder is type II diabetes.
- (Original) A method according to Claim 1 wherein the disorder is Creutzfeldt-Jakob disease.
- 27. (Original) A method according to Claim 1 wherein the disorder is a Huntington's disease.
- 28. (Original) A method according to Claim 1 wherein the disorder is macular degeneration.
- 29. (Original) A method according to Claim 1 wherein the disorder is a prion disease.
- 30. (Original) A method according to Claim 1 wherein Step A comprises:

obtaining sample cells from the subject;

extracting DNA from the sample cells;

subjecting the extracted DNA to mitochondrial DNA control region amplification;

determining whether homoplasmic 414 and 477 nucleotide variants are present by direct sequencing for heteroplasmic 414 and 477 nucleotide mutations; and

if 414 and 477 nucleotide variants are detected, cloning the mutant molecules and sequencing the clone.

- 31. (Cancelled)
- 32. (Cancelled)
- 33. (Cancelled)
- 34. (Cancelled)